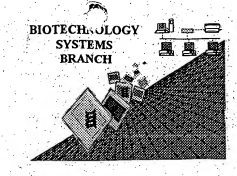
RAW SEQUENCE LISTING ERROR REPORT



S7-57-16 ₩V

The Biotechnology Systems Branch of the Scientific and Technical Information Center (STIC) detected errors when processing the following computer readable form:

Application Serial Number: 09/753,892Source: 0/PEDate Processed by STIC: 1/23/2001

THE ATTACHED PRINTOUT EXPLAINS DETECTED ERRORS.
PLEASE FORWARD THIS INFORMATION TO THE APPLICANT BY EITHER:

1) INCLUDING A COPY OF THIS PRINTOUT IN YOUR NEXT COMMUNICATION TO THE APPLICANT, WITH A NOTICE TO COMPLY or,

2) TELEPHONING APPLICANT AND FAXING A COPY OF THIS PRINTOUT, WITH A NOTICE TO COMPLY

FOR CRF SUBMISSION QUESTIONS, PLEASE CONTACT MARK SPENCER, 703-308-4212.

FOR SEQUENCE RULES INTERPRETATION, PLEASE CONTACT ROBERT WAX, 703-308-4216. PATENTIN 2.1 e-mail help: patin21help@uspto.gov or phone 703-306-4119 (R. Wax) PATENTIN 3.0 e-mail help: patin30help@uspto.gov or phone 703-306-4119 (R. Wax)

TO REDUCE ERRORED SEQUENCE LISTINGS, PLEASE USE THE CHECKER VERSION 3.0 PROGRAM, ACCESSIBLE THROUGH THE U.S. PATENT AND TRADEMARK OFFICE WEBSITE. SEE BELOW:

Checker Version 3.0

The Checker Version 3.0 application is a state-of the-art Windows based software program employing a logical and intuitive user-interface to check whether a sequence listing is in compliance with format and content rules. Checker Version 3.0 works for sequence listings generated for the original version of 37 CFR §§1.821 – 1.825 effective October 1, 1990 (old rules) and the revised version (new rules) effective July 1, 1998 as well as World Intellectual Property Organization (WIPO) Standard ST.25

Checker Version 3.0 replaces the previous DOS-based version of Checker, and is Y2K-compliant. Checker allows public users to check sequence listings in Computer Readable form (CRF) before submitting them to the United States Patent and Trademark Office (USPTO). Use of Checker prior to filing the sequence listing is expected to result in fewer errored sequence listings, thus saving time and money

Checker Version 3.0 can be down loaded from the USPTO website at the following address: http://www.uspto.gov/web/offices/pac/checker

	ERROR DETECTED	SUGGESTED	CORRECTION	;	SERIAL NUMBER: _	09/753,892	
ATTN	I: NEW RULES CASES: PI Wrapped Nucleics	The number/text a	at the end of each line "	'wrapped" down to t	he next line.	D BY PTO SOFTWARE	
			your file was retrieved in the right margin to .3, as				
2	Wrapped Aminos	This may occur if	umber/text at the end of f your file was retrieved or right margin to .3, as	in a word processo	r after creating it.	ne.	
3	Incorrect Line Length	The rules require	that a line not exceed 7	 2 characters in leng	yth. This includes spa	aces.	
4	Misaligned Amino Acid Numbering		nder each 5th amino ac pering. It is recommend				
5	Non-ASCII		aved in ASCII (DOS) to ur subsequent submiss			e processed.	
6	Variable Length	As per the rules, e	contain n's or Xaa's weach n or Xaa can only of maximum number of of the feature section that so	represent a single ro each residue having	esidue. variable length and		
7	PatentIn ver. 2.0 "bug"	sequence(s) previously coded to the subsequent	n version 2.0 has cause Normally, F nucleic acid sequence. t amino acid sequence. ficial or Unknown seq	Patentin would auto Please manually c This applies prin	matically generate this opy the relevant <220:	s section from the)>-<223> section	
8	Skipped Sequences (OLD RULES)	(2) INFORMATION (i) SEQUENCE CONTROL (xi) SEQUENCE (missing. If intentional, N FOR SEQ ID NO:X: HARACTERISTICS:(COESCRIPTION:SEQ ID intentionally skipped)	Do not insert any hea D NO:X:		skipped sequence: ENCE CHARACTERISTICS	")
		Please also adjust	the "(iii) NUMBER OF	SEQUENCES:" re	sponse to include the	skipped sequence(s).	
9	Skipped Sequences (NEW RULES)	Sequence(s) <210> sequence <400> sequence 000		please use the follo	wing format for each s	skipped sequence.	
0	Use of n's or Xaa's (NEW RULES)	Use of <220> to <	Kaa's have been delecto 223> is MANDATORY section, please explain	if n's or Xaa's are p	resent.	n or Xaa represents.	
1	Use of <213>Organism (NEW RULES)	Sequence(s)	are missing this m	andatory field or its	response.		
2	Use of <220>Feature (NEW RULES)	Please explain so	are missing the <220> 223> is MANDATORY ource of genetic mate Register," 6/01/98	if <213>ORGANIS rial in <220> to <2	M is "Artificial" or "Uni 23> section.		ules)
3	Patentln ver. 2.0 "bug"		e "Copy to Disk" func ssing mandatory numer			ses a corrupted d on raw sequence listing).	

Instead, please use "File Manager" or any other means to copy file to floppy disk.

OIPE

RAW SEQUENCE LISTING

PATENT APPLICATION: US/09/753,892

Output Set: N:\CRF3\01232001\1753892.raw

Input Set : A:\pana-0002.txt

DATE: 01/23/2001

TIME: 10:41:48

Does Not Comply
Corrected Diskette Needed

3 <110> APPLICANT: Yakubov, Leonid 5 <120> TITLE OF INVENTION: COMPOSITIONS COMPRISING GENOME SEGMENTS AND METHODS OF USING THE SAME 7 <130> FILE REFERENCE: PANA-0002 delete servicion 9 <140> CURRENT APPLICATION NUMBER: US/09/753,892 9 <141> CURRENT FILING DATE: 2001-01-03 9 <160> NUMBER OF SEQ ID NOS: 1 11 <170> SOFTWARE: PatentIn version 3.0 13 <210> SEQ ID NO: 1 14 <211> LENGTH: 22 ation for Artificial Sequence or Unknown 15 <212> TYPE: DNA 16 <213> ORGANISM: (Artificial/Unknown) 18 <220> FEATURE: 19 <221> NAME/KEY: misc_feature goes on (2237 20 <222> LOCATION: (11)..(16) 21 <223> OTHER INFORMATION: N is any or unknown Pline. Do not 24 < TUO> SEQUENCE: 1 *42*237 > 25 ccgactcgag nnnnnnatgt gg insert it on 12137 line.

Per new Sequence Rules, the only valid 22137 remonses

are: Unknown, Artificial Sequence, or scientific name (Genus/species),

one of the three -do not combine responses, as

shown about. See circled portion of them 12 on Evon

Sunnay Sheet.

VERIFICATION SUMMARY

DATE: 01/23/2001 TIME: 10:41:49

PATENT APPLICATION: US/09/753,892

Input Set : A:\pana-0002.txt
Output Set: N:\CRF3\01232001\1753892.raw

L:9 M:270 C: Current Application Number differs, Replaced Current Application No L:9 M:271 C: Current Filing Date differs, Replaced Current Filing Date L:25 M:341 W: (46) "n" or "Xaa" used, for SEQ ID#:1